

certain characteristics of the antibody, acute renal failure, and infection.

There are few situations in hematology where one laboratory test is as informative as the direct antiglobulin test developed in 1945 by R. R. A. Coombs and known since then by this name. The currently recommended version of this test uses in the first instance a “broad-spectrum” reagent, i.e., one that will detect not only immunoglobulins (Ig) but also complement (C) components (usually C3 fragments) bound to the surface of the patient’s red cells. If the test is positive (and barring special circumstances such as previous blood transfusion), it is practically diagnostic of AIHA, and one can then determine, by using specific reagents, whether Ig or C or both are implicated. The sensitivity of the Coombs test varies depending on the techniques that are used: in general, the test is positive if there are an average of at least 400 molecules of Ig and/or C on each red cell; but with more advanced techniques involving flow cytometry analysis or enzyme-linked radiolabeled tests allowing the detection of ~30–40 antibody molecules per erythrocyte, the sensitivity can be pushed to as low as 30–40 molecules per red cell. Therefore liaison with a specialized laboratory is desirable; a dual direct antiglobulin test has also been developed. In the past the diagnosis of “Coombs-negative AIHA” was regarded as a last resort, but it is important to know that a patient with this label may have severe AIHA, because if the antibody is powerful (high affinity/avidity), few molecules may be sufficient to opsonize red cells. Based on the Coombs test findings as well as on the thermal characteristics and the antigenic specificities of the autoantibodies ([Table 100-7](#)), AIHA has been classified into subtypes.

TABLE 100-7 Classification of Acquired Immune Hemolytic Anemias